Bioinformatic approaches to regulatory genomics and epigenomics

376-1347-00L - 2023 | week 02

Pierre-Luc Germain



Plan for today

Debriefing on the assignments

- The notion of gene
- Genome builds and transcriptome assemblies

- Practical:
 - AnnotationHub and EnsDb objects
 - GenomicRanges and their manipulations

Debriefing on the assignments

```
#install.packages(c("gsl","ggplot2"))
# I already installed Bioconductor

#BiocManager::install(c("GenomicRanges", "rtracklayer", "EnrichedHeatmap", "AnnotationHub", "ensembldb", "edgeR", "esATAC", "sechm","genomation","Rsubread","Rfastp"))
#install.packages("cowplot")
#BiocManager::install("ETHZ-INS/epiwraps")
```

"I decided to comment the install commands so that R does not install those packages everytime I ran the code since it is only necessary once"

In such circumstances, you can set whole chunks not to be run:

""{r, eval=FALSE} some_code() # this code chunk will be displayed in the document, but won't be run to be run:

Debriefing on the assignments

Include all the code chunks which produce output:

```
fr pressure, echo=FALSE}
sessionInfo()

info
info
sessionInfo()

info
sessionInfo()
```

... or why nobody knows how many genes we have

1900 - Rediscovery of Mendel's work (1860s) 1903 - Chromosomes are hereditary units 1913 - Chromosomes are linear arrays of genes 1941 - Beadle & Tatum: the one-gene-one-enzyme hypothesis 1944 - DNA is the genetic material 1951 - First protein sequences 1953 - DNA is a double helix 1961 - Jacob and Monod: the *lac* operon 1977 - DNA sequencing 1977 - Eukaryotic genes are spliced 1995 - First bacterial genomes sequenced 2000 - Next Generation Sequencing (NGS) 2001 - Draft of the human genome 2003 - RNA-seq 2006 - ChIP-seq 2012 - ENCODE, ATAC-seq

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1944 - DNA is the genetic material

1951 - First protein sequenced

1953 - DNA is a double helix

1961 - Jacob and Monod: the *lac* operon

1977 - DNA sequencing

1977 - Eukaryotic genes are spliced

1995 - First bacterial genomes sequenced

2000 - Next Generation Sequencing (NGS)

2001 - Draft of the human genome

2003 - RNA-seq

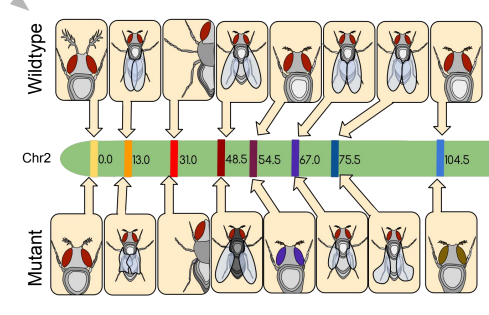
2006 - ChIP-seq

2012 - ENCODE, ATAC-seq

Classical genetics:

A "gene" is a unit of

heredity



(Morgan's Drosophila genetic map, adapted from Twaanders17, CC BY-SA 4.0, via Wikimedia Commons)

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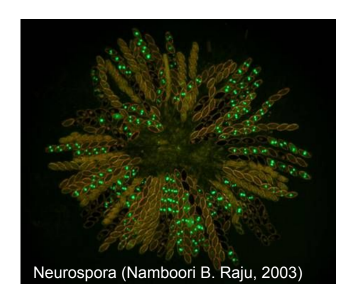
Classical genetics:

Early molecular genetics:

A "gene" is a unit of heredity

(+)

A "gene" is a part of DNA that encodes for a protein



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Classical A "gene" is a unit of genetics: heredity

Early A "gene" is a part of molecular DNA that encodes genetics: for a protein

- "structural" genes encode for proteins
- "regulatory" genes → regulate the structural genes

J. Mol. Biol. (1961) 3, 318-356

REVIEW ARTICLE

Genetic Regulatory Mechanisms in the Synthesis of Proteins†

François Jacob and Jacques Monod

Services de Génétique Microbienne et de Biochimie Cellulaire, Institut Pasteur, Paris

(Received 28 December 1960)

The synthesis of enzymes in bacteria follows a double genetic control. The so-called structural genes determine the molecular organization of the proteins. Other, functionally specialized, genetic determinants, called regulator and operator genes, control the rate of protein synthesis through the intermediacy of cytoplasmic components or repressors. The repressors can be either inactivated (induction) or activated (repression) by certain specific metabolites. This system of regulation appears to operate directly at the level of the synthesis by the gene of a short-lived intermediate, or messenger, which becomes associated with the ribosomes where protein synthesis takes place.

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Classical A "gene" is a unit of genetics: heredity

Early A "gene" is a part of molecular DNA that encodes genetics: for a protein

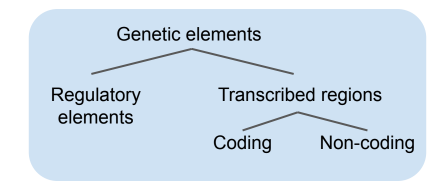
- "structural" genes encode for proteins
- "regulatory" genes → regulate the structural genes

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Nano-lectures:

- A 24sec technical summary
- A 7-words summary everyone can understand

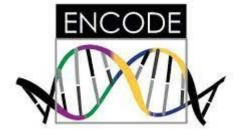
So what's a gene today?

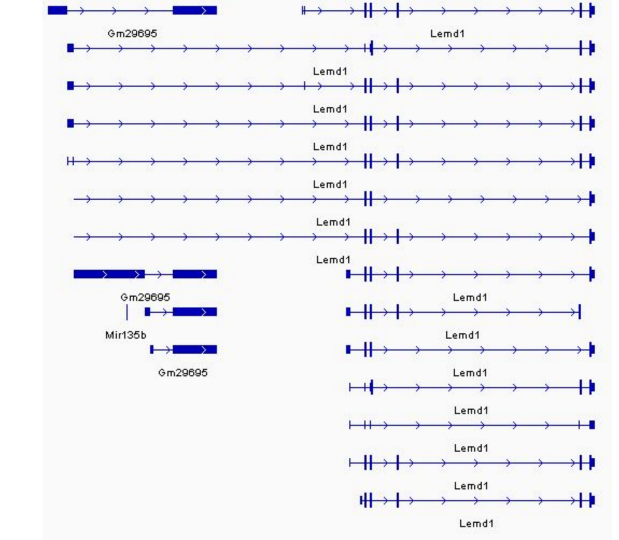
"The gene is a union of genomic sequences encoding a coherent set of potentially overlapping functional products"

(Gerstein et al., 2007)

"On this view, genes represent a higher-order framework around which individual transcripts coalesce, creating a poly-functional entity that assumes different forms under different cellular states, guided by differential utilization of regulatory DNA."

(Stamatoyannopoulos, 2012)





Transcripts tend to

genes depending

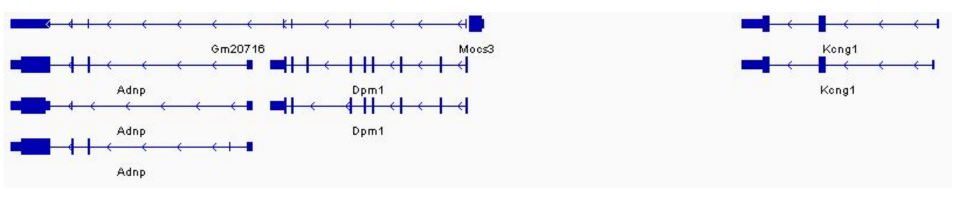
overlapping exonic

be grouped by

on partially

sequences

But of course that's not a universal rule...



Reference genomes and gene annotations

The **reference genome** refers to the **sequence** of a genome.

For mainstream organisms (e.g. human/mouse), there is one standardized genome (from the Genome Reference Consortium) which everyone uses, but different **build** versions of that genome, e.g.:

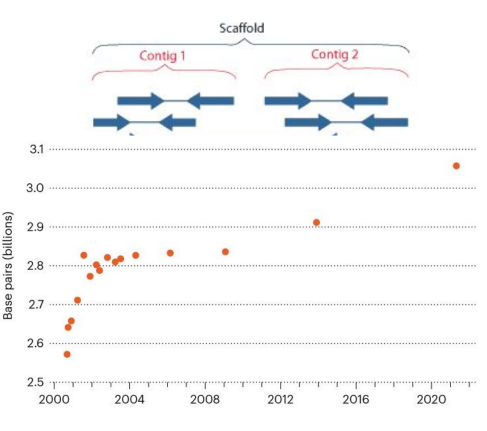
Human:

- o GRCh38/hg38 (released in 2013)
- o GRCh37/hg19 (released in 2009)
- 0 ...

Mouse:

- o GRCm39/mm39 (released in 2020)
- o GRCm38/mm10 (released in 2011)
- GRCm37/mm9 (released in 2007)

0 ...



(Reardon, Nature News 2021)

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Within each genome build, there are also patches:

Human:

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- GRCh37/hg19 (released in 2009)
- 0 ..

Mouse:

- GRCm39/mm39 (released in 2020)
- o GRCm38/mm10 (released in 2011)
- o GRCm37/mm9 (released in 2007)

o ..

GRCh38.p13 (released in 2019) GRCh38.p12 (released in 2017)

Coordinates are stable within a build, but not across builds

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- o ...

Annotations refer to the catalogues of regulatory elements, genes and transcripts associated to a genome.

There are two main sources for gene annotations (aka "gene builds"):

Ensembl

104 (may 2021)

103 (nov 2020)

102 (april 2020)



example transcript:

ENST000012343.2

ENST000012343.1

ENST000012343.1



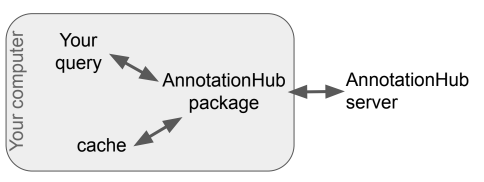
Accessing ensembl annotations

http://ftp.ensembl.org/pub/

- contains various types of data, most importantly:
 - .fasta files: sequences
 - (e.g. DNA or cDNA)
 - o .gtf/.gff files: gene models
 - (i.e. exon coordinates and inclusion into transcripts)

AnnotationHub

Standardized access to a large variety of annotations



including genomes, gene annotations (e.g. EnsDb objects) and more

Further information

See the documentation of the <u>ensembldb</u> package for manipulating EnsDb objects

 See the documentation of the <u>GenomicRanges</u> package for manipulating GRanges objects (and their derivatives)

Assignment for this week

- Using AnnotationHub, find and download the following annotations data:
 - The mouse (Mus Musculus) EnsDb object, version 102, genome build GRCm38
 - The mouse genome sequence (dna sm) in TwoBit/2bit format for GRCm38
 - The drosophila melanogaster genome sequence (dna_sm) in TwoBit/2bit format for BDGP6

- Using the mouse EnsDb, find the following:
 - How many different *ensembl gene IDs* and *gene symbols* are there for protein-coding genes?
 - Plot the distribution of the (spliced) length of protein-coding transcripts
 - (tip: this will require you to extract exons of protein-coding transcripts from the database, and split them by transcript, before summing the width of the exons of each transcript)

Name your markdown file 'assignment.Rmd', render it, and put it (along with the produced html) in the week02 folder of your repository, and push!